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Deep Blue Oxadiazole-Containing Thermally Activated Delayed Fluorescence Emitters For Organic Light-Emitting Diodes

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Abstract

A series of four novel deep blue to sky blue thermally activated delayed fluorescence (TADF) emitters (**2CzdOXDMe**, **2CzdOXD4MeOPh**, **2CzdOXDPh** and **2CzdOXD4CF₃Ph**) have been synthesized and characterized. These oxadiazole-based emitters demonstrated bluer emission compared with reference emitter **2CzPN** thanks to the

weaker acceptor strength of oxadiazole moieties. The oxadiazole compounds doped in hosts (mCP and PPT) emitted from 435-474 nm with photoluminescence quantum yields ranging from 14%-55%. The emitters possess singlet-triplet excited state energy gaps (ΔE_{ST}) between 0.25 and 0.46 eV resulting in delayed components ranging from 4.8 to 25.8 ms. The OLED device with **2CzdOXD4CF₃Ph** shows a maximum external quantum efficiency of 11.2% with a sky-blue emission at CIE of (0.17, 0.25) while the device with **2CzdOXD4MeOPh** shows a maximum external quantum efficiency of 6.6% with a deep-blue emission at CIE of (0.15, 0.11).

Introduction

In organic light-emitting diodes (OLEDs), light is produced through radiative decay of electrically generated excitons. As a function of spin statistics, 75% of the excitons are triplets and 25% of the excitons are singlets. In order to obtain efficient electroluminescent (EL) devices, 100% of the excitons need to be harvested within the emissive layer of the device.¹⁻² The current state-of-the-art emitters for OLEDs rely on neutral iridium(III) and platinum(II) complexes as, (1) the emission color of these materials can be easily tuned as a function of ligand design, (2) these complexes possess microsecond emission lifetimes and, most importantly, (3) the heavy metal mediates an efficient intersystem crossing (ISC) to harness both singlets and triplets in OLED devices.³⁻⁷ Despite internal quantum efficiencies (IQE)⁵ approaching 100%, these organometallic emissive materials are costly and toxic and, importantly, there exists no stable and bright deep blue emitter, a key component for both displays and lighting applications.⁸⁻⁹

As a response to the drawbacks of phosphorescent organometallic emitters, thermally activated delayed fluorescent (TADF)¹⁰⁻¹⁴ materials have recently come to the fore, as they too are capable of recruiting 100% of the excitons because the energy gap between the lowest singlet (S_1) and triplet (T_1) levels, (ΔE_{ST}), is small enough to permit reverse intersystem crossing (RISC) between the two states at ambient temperature.¹⁵ Compounds possessing small ΔE_{ST} possess structures wherein the highest occupied molecular orbital (HOMO) is spatially separated from the lowest unoccupied molecular orbital (LUMO) in order to minimize the exchange integral between the two.¹⁵⁻¹⁷ Importantly, most TADF emitters are based on either cheap complexes mostly derived from copper¹⁸⁻²⁴ or are purely organic in nature.^{15,25-29}

One of the grand challenges remaining in emitter design for OLEDs is the development of stable and bright *deep-blue* emitters (CIE x,y coordinates each < 0.2),^{8,14,30-33} as only limited examples of *deep-blue* TADF emitters have been reported to date. The first organic deep-blue TADF emitters, reported by Zhang *et al.* in 2012, were based on a diphenylsulfone (DPS) acceptor.¹⁶ OLEDs using these emitters showed EQE_{max} of 10% and CIE at (0.15, 0.07). Since this report, other analogues based on DPS acceptors have been reported, many reaching EQE_{max} values in excess of 10%.³⁴⁻³⁸ Employing a related TADF emitter based on a dibenzo-fused phosphacycle acceptor produced a somewhat poorer performance in the device, with EQE_{max} of 4.9% and CIE at (0.15, 0.16).³⁹ Hatakeyama *et al.*⁴⁰ designed an interesting class of deep-blue TADF emitters based on a “multiple resonance effect”, which produced narrow-band emission spectra with CIE at (0.13, 0.09) and an EQE_{max} of 13.5%. A second generation emitter by the same group showed improved performance with EQE_{max} of 18.3% and CIE at (0.13, 0.11).⁴¹ Cui *et al.*⁴² synthesized a series of deep-blue TADF emitters based on a carbazole donor and a diphenyltriazine acceptor and

achieved an EQE_{max} of 19.2% with CIE at (0.15, 0.10). Wada *et al.*⁴³ adapted an earlier emitter design of Hirata *et al.*²⁹ by the incorporation of adamantyl groups on the triazine acceptor to afford deep-blue TADF emitter for solution-processed OLEDs, showing EQE_{max} of 11.2% and CIE at (0.15, 0.13) while Woo *et al.*⁴⁴ have modified the donor groups to both increase emission energy, as well as achieving preferential horizontal transition dipole orientation of the emitters in the film in order to enhance light outcoupling; achieving devices with EQE_{max} of 7.7% and CIE at (0.15, 0.08). Komatsu *et al.*⁴⁵ developed related pyrimidine-based emitters with EQE_{max} of 18% and CIE at (0.16, 0.15). Seo *et al.*⁴⁶ employed an indolocarbazole moiety as the acceptor in the design of their deep-blue TADF emitters, which achieved an EQE of 19.5% with CIE at (0.15, 0.16). Lee *et al.*⁴⁷ demonstrated that by placing a carbazole donor *ortho* to a dimesitylborane acceptor, a very high EQE_{max} of 22.6% with CIE of (0.14, 0.15) could be achieved. When both orientation in the film and photoluminescence quantum yield can be optimized, outstanding device performance can result, as demonstrated by Rajamalli *et al.*,⁴⁸ who produced OLEDs with EQE_{max} of 31.9% and CIE of (0.14, 0.18) based on a dipyritylketone acceptor design. Recently, a deep-blue emitter bearing two carbazole donors *ortho* to the cyanobenzene acceptor was reported by Chan *et al.*,⁴⁹ where OLEDs achieved EQE_{max} of 10.3% and CIE at (0.16, 0.06). The device performances of the aforementioned work together with this study are summarized in Table 1.

Table 1. Summary of reported deep-blue TADF device performances and this work.

Emitter	EQE _{max} (%)	EQE @ 100 cd m ⁻² (%)	CIE	Ref.
2CzdOXD4MeOPh	6.6	2.0	(0.15, 0.11)	This work
3	10	~2.5	(0.15, 0.07)	16
DMOC-DPS	14.5	9.0	(0.16, 0.16)	34
DTC-mBPSB	5.5	---	(0.15, 0.08)	35
DMTDAc	19.8	19.8	(0.15, 0.13)	36
DTPPDDA	4.7	2.2	(0.15, 0.09)	37

G2	--- (4.1 cd A ⁻¹) ^a	---	(0.15, 0.12)	38
MFAc-OPO	4.9	~0.6	(0.15, 0.16)	39
DABNA-1	13.5	~6	(0.13, 0.09)	40
B2	18.3	12.6	(0.13, 0.11)	41
Cz-TRZ3	19.2	---	(0.15, 0.10)	42
FA-TA	11.2	---	(0.15, 0.13)	43
DTXSAF	7.7	~5	(0.15, 0.08)	44
Ac-3MHPM	18	10.4	(0.16, 0.15)	45
ICzDAc	19.5	---	(0.15, 0.16)	46
CzoB	22.6	18.4	(0.14, 0.15)	47
3DPyM-pDTC	31.9	26.1	(0.14, 0.18)	48
DCzBN3	10.3	5.4	(0.16, 0.06)	49
^a EQE not reported.				

In this study, a series of four deep blue to sky blue TADF emitters have been synthesized and characterized whose structures are shown in Chart 1. These molecules possess the same scaffold as the dicarbazoyldicyanobenzene (**2CzPN**) reported by Uoyama *et al.*¹⁵ but with the replacement of the cyano groups by the less electron-withdrawing oxadiazole units. The rationale for our design is three-fold. Firstly, the weaker acceptor strength of oxadiazoles⁵⁰⁻⁵¹ compared with cyano groups will translate into a bluer emission, presuming that the intramolecular charge transfer (ICT) nature of emission is conserved. Secondly, oxadiazoles show promising thermal stability and electron injection and transporting properties.^{50,52-57} Lastly, according to theoretical calculations, the LUMO density of **2CzPN** is mostly located on the central benzene ring¹⁵ whereas that of oxadiazole is located on the heterocyclic ring (*vide infra*).^{50,54} This implies that the electron density of the HOMO, which in this study is localized on the carbazole units, and LUMO in the oxadiazole derivatives should be more separated than in **2CzPN**; however, the weaker strength of the

oxadiazole acceptor results in greater delocalization of the electron density of the LUMO onto the benzene bridge thus resulting in comparable ΔE_{ST} to the reference emitter.

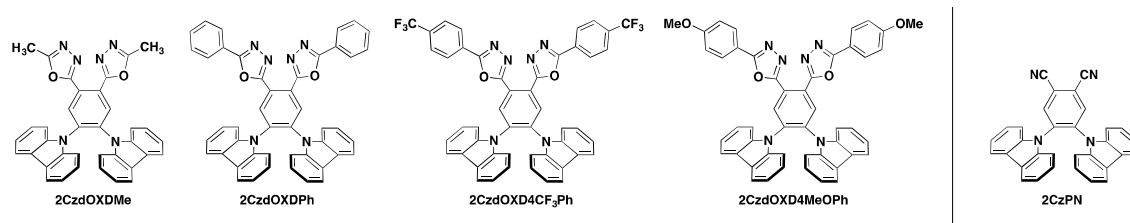
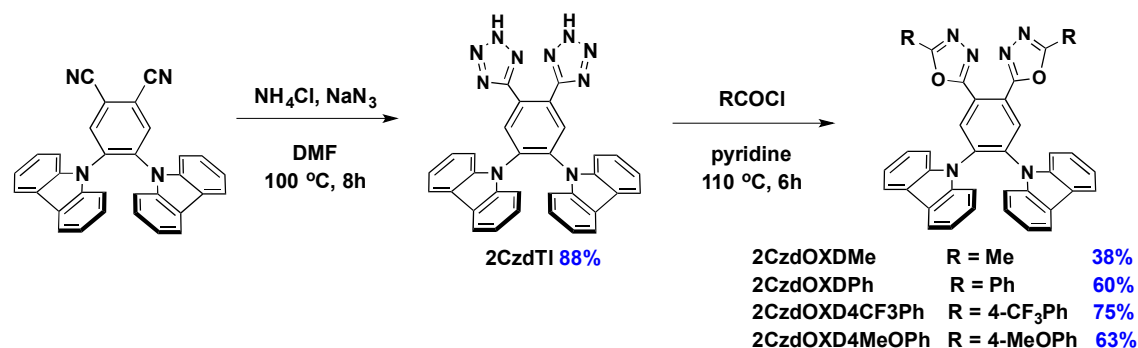


Chart 1. Left: Oxadiazole TADF emitters in this study. Right: Reference emitter **2CzPN**

Results and Discussion

Synthesis



Scheme 1. Synthesis of oxadiazole-based emitters.

The synthetic route for the four oxadiazole emitters is shown in Scheme 1, starting from the reference emitter **2CzPN**, which is itself obtained in a much improved yield of 80% compared to the one originally reported by Uoyama *et al.* (8.5%) using a similar protocol.¹⁵ Despite the many synthetic routes available for the installation of the oxadiazole moiety,⁵⁸⁻⁶⁰ a two-step protocol passing through a ditetrazole intermediate (**2CzdTI**)⁵⁹ was found to be most straightforward given the presence of the cyano groups. As a result, **2CzPN** was reacted with ammonium chloride and NaN₃ in DMF at 120 °C overnight to afford **2CzdTI** in 88% yield. The key for the success of this reaction is the isolation of **2CzdTI**, which can be

precipitated from the reaction mixture as a white solid by pouring the reaction mixture into a 1M HCl solution. The ease of isolation of **2CzdTI** permits facile multigram scale up of the target materials. Following isolation, **2CzdTI** can then be reacted with the corresponding acid chloride of choice to provide the desired oxadiazole emitters in good yield (38-75%). All four oxadiazole emitters were found to be thermally stable beyond 300 °C by TGA; **2CzdOXDMe** and **2CzdOXDPh** melt at 308 and 310 °C, respectively while **2CzdOXD4CF₃Ph** and **2CzdOXD4MeOPh** melt at 274 and 260 °C, respectively (For TGA, DTA and DSC traces see: Figures S28-S39).

Absorption and Electrochemistry

Table 2. Summary of absorption and electrochemistry of oxadiazole emitters and **2CzPN** as control.

Emitter	λ_{abs}^a (nm), [$\epsilon (\times 10^4 \text{ M}^{-1} \text{ cm}^{-1})$]	Electrochemistry ^b (eV)
2CzdOXDMe	281 [2.26], 290 [2.46], 321 [1.37], 335 [1.51], 353 [1.36]	HOMO: -5.83 LUMO: -2.70 ΔE : 3.13
2CzdOXDPh	258 [5.38], 283 [4.39], 291 [4.48], 320 [2.38], 333 [2.20], 361 [2.12]	HOMO: -5.84 LUMO: -2.80 ΔE : 3.04
2CzdOXD4CF₃Ph	257 [4.81], 283 [4.36], 291 [4.45], 319 [2.00], 332 [1.95], 367 [1.79]	HOMO: -5.84 LUMO: -2.86 ΔE : 2.98
2CzdOXD4MeOPh	256 [4.10], 290 [4.72], 334 [2.45], 353 [2.12]	HOMO: -5.88 LUMO: -2.74 ΔE : 3.14
2CzPN	280(sh) [1.72], 289 [2.14], 319 [1.02], 329 [1.17], 364 [1.14] ^c	HOMO: -5.88 LUMO: -2.97 ΔE : 2.91
2CzPN^d	367 ^e	HOMO: -5.89 LUMO: -2.97 ΔE : 2.92

^a in DCM at 298 K. ^b in MeCN with 0.1 M [*n*Bu₄N]PF₆ as the supporting electrolyte and Fc/Fc⁺ as the internal reference. The HOMO and LUMO energies were calculated using the relation $E_{\text{HOMO/LUMO}} = -(E_{\text{pa,1}}^{\text{ox}}/E_{\text{pc,1}}^{\text{red}} + 4.8)\text{eV}$, where $E_{\text{pa}}^{\text{ox}}$ and $E_{\text{pc}}^{\text{red}}$ are anodic and cathodic peak potentials respectively. $\Delta E = -(E_{\text{HOMO}} - E_{\text{LUMO}})$.⁶¹ ^c in MeCN at 298K. ^d Values from Ref⁶². ^e Only longest wavelength peak reported, in MeCN.

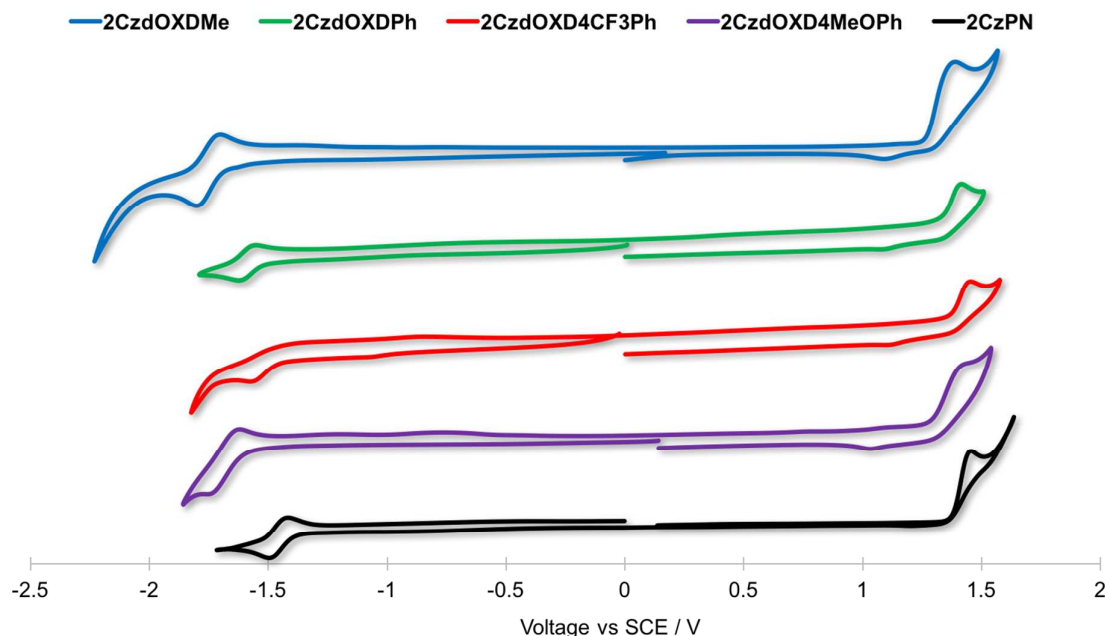


Figure 1. CVs of the TADF emitters in degassed MeCN under argon with 0.1 M $[n\text{Bu}_4\text{N}]\text{PF}_6$ as the supporting electrolyte and using Fc/Fc^+ as the internal standard. The CVs are reported vs SCE.

The electrochemistry of the TADF emitters was studied by CV in degassed MeCN solutions under argon, Figure 1. The HOMO levels are expectedly nearly identical (Table 2, -5.83 to -5.88 eV), the result of the very similar orbital delocalization patterns over the carbazole donors and the central phenyl bridge as shown by Density Functional Theory (DFT) calculations (*vide infra*). The LUMO levels of the different emitters are, however, strongly influenced by the acceptor strength induced by the groups attached to the oxadiazole acceptors. The LUMO level of **2CzdOXDPh** is lower by 0.1 eV compared to **2CzdOXDMe** as a result of increased conjugation length afforded by the phenyl group. Compound **2CzdOXD4CF₃Ph** has the most stabilized LUMO (-2.86 eV) due to the strong electron-withdrawing effect of the trifluoromethyl group while **2CzdOXD4MeOPh** has the shallowest LUMO (-2.74 eV) due to the strong mesomerically electron-donating effect of methoxy group. For each emitter, the oxidation is irreversible, which is not unexpected as carbazole

radical cations are known to be electrochemically unstable and undergo dimerization.⁶³⁻⁶⁴ Conversely, only **2CzdOXD4CF₃Ph** shows an irreversible reduction, probably due to cleavage of the C-F bonds.⁶⁵⁻⁶⁶ The result of the use of the oxadiazole acceptors is an increase in the electrochemical gap from 2.91 eV for **2CzPN** to between 2.98-3.14 eV for the four TADF emitters.

Photophysical Properties

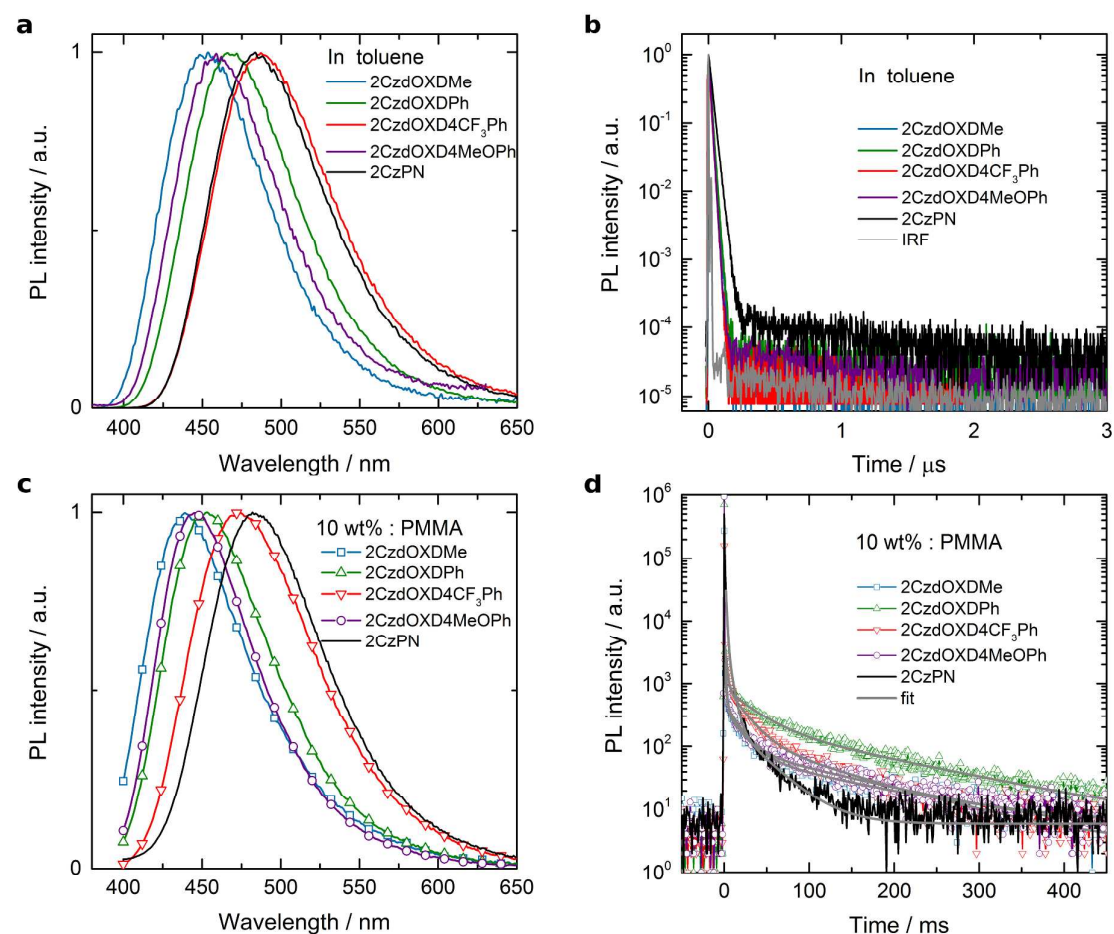


Figure 2. a) Emission spectra ($\lambda_{\text{exc}} = 340$ nm) and b) decay profiles ($\lambda_{\text{exc}} = 378$ nm) of the TADF emitters in degassed toluene at RT; c) Steady-state photoluminescence spectra of emitters embedded in a PMMA matrix at 10 wt% ($\lambda_{\text{exc}} = 378$ nm); d) Photoluminescence decay of emitters embedded in PMMA matrix at 10 wt% ($\lambda_{\text{exc}} = 378$ nm).

Table 3. Summary of solution and thin film photophysical properties of the oxadiazole TADF emitters and **2CzPN** as reference.

	PhMe			Thin Film ^c		
	λ_{PL}^a /nm	Φ_{PL}^b /%	τ_{PL} /ns	λ_{PL}^a /nm	Φ_{PL}^d /%	τ_{PL} /ns ^e
2CzdOXDMe	453 (79)	28.7 (24.5)	14.5	439 (77)	46.4 (39.6)	11.8, $3.0 \cdot 10^6$ (75%), $40.0 \cdot 10^6$ (19%)
2CzdOXDPh	466 (82)	38.3 (27.5)	15.6	453 (81)	62.0 (54.5)	11.0, $4.9 \cdot 10^6$ (63%), $45.2 \cdot 10^6$ (27%)
2CzdOXD4CF₃Ph	487 (90)	39.1 (25.7)	19.1	474 (94)	74.9 (57.1)	11.6, $2.3 \cdot 10^6$ (77%), $23.2 \cdot 10^6$ (19%)
2CzdOXD4MeOPh	459 (78)	47 (46.7)	13.1	447 (73)	41.1 (37.7)	11.5, $1.5 \cdot 10^6$ (83%), $17.3 \cdot 10^6$ (10%)
2CzPN	484 (85)	28.1 (26.2)	24.4 (99.5%)/1122 (0.5%)	492 (83)	76.0 (62.6)	18.4, $1.2 \cdot 10^6$ (90%), $4.8 \cdot 10^6$ (10%)

^a Emission maxima and full-width at half maximum (FWHM) are reported from degassed solutions. FWHM in parentheses. ^b 0.5M quinine sulfate in H₂SO₄ (aq) was used as reference (PLQY: 54.6%).⁶⁷ Values quoted are in degassed solutions. Values in parentheses are for aerated solutions. ^c Thin films were prepared by spin-coating doped samples in PMMA (10 wt%). ^d Values determined using an integrating sphere under an N₂ atmosphere as described above in the main text. Values in parentheses are for samples measured in air. ^e Prompt lifetimes determined from the monoexponential fit of the initial decay by TCSPC; delayed lifetimes determined from the bi-exponential decay by MCS, weighting corresponding to the pre-exponential factors are indicated in parentheses.

The solution-state photophysical properties of the four oxadiazole emitters were studied in toluene (Figure 2a), DCM and MeCN. The photophysical data for PhMe are reported in Table 3. All four emitters show positive solvatochromism and broad and unstructured emission profiles,^{59,68-69} which are characteristic of emission from an ICT state. The broadness of emission also increases with increasing polarity of the solvent. Regardless of solvent, the wavelength at emission maximum increases in the order: **2CzdOXDMe** < **2CzdOXD4MeOPh** < **2CzdOXDPh** < **2CzdOXD4CF₃Ph** (Figure S41), which is consistent with the correspondingly decreasing bandgaps obtained from electrochemistry.

We envisioned that the weaker acceptor strength of the oxadiazole compared to the cyano group would cause a desired blue-shift in the emission in our emitters with respect to **2CzPN**. Indeed, the LUMO levels of the four oxadiazole compounds range from -2.70 eV to -2.86 eV while the LUMO of **2CzPN** was determined to be -2.97 eV (Table 2), and their emission was blue-shifted as a consequence. For example, **2CzdOXDMe** has an emission maximum at 453 nm in toluene, blue-shifted by 31 nm (1414 cm⁻¹) compared with **2CzPN** (484 nm) in the same solvent.

The photoluminescence quantum yields (Φ_{PL}) in degassed solutions range from 28 to 47% and do not significantly change with solvent choice. The Φ_{PL} values in degassed solutions are always higher than those in aerated solutions. This finding provides direct evidence for transient population of the triplet excited state, which is sensitive to triplet oxygen quenching and is diagnostic of TADF materials.⁶⁹⁻⁷² The emission lifetime of **2CzPN** in toluene consists of a prompt (24.4 ns) and a weak delayed component (1.12 μ s, 0.5% amplitude in multi-exponential fit). The prompt component in the oxadiazole-containing compounds in toluene

is faster (13-19 ns); however, the delayed component is too weak to be resolved as it coincides with the instrument response function (Figure 2b, grey line). Thus, the delayed fluorescence in all of the studied compounds in toluene is smaller than we can reliably detect.

To assess the viability of these oxadiazole emitters in OLED devices, their photophysical properties were investigated in the solid state. Thin films were prepared by spin-coating 10 wt % DCM solution of emitter in PMMA (Figure 2c). All emission maxima are blue-shifted by ~15 nm and the profiles generally slightly sharper compared with those measured in toluene solution. The thin film Φ_{PL} (40-75%) are significantly higher than in solution as a result of the more rigid environment that constrains the torsional motion thereby reducing the non-radiative decay rate.⁷³ In particular, **2CzdOXDPh** ($\lambda_{\text{PL}} = 453 \text{ nm}$) and **2CzdOXD4CF₃Ph** ($\lambda_{\text{PL}} = 474 \text{ nm}$) exhibit remarkable Φ_{PL} values of 62% and 75% in the deep-blue region under nitrogen, making them very promising blue TADF materials for OLEDs. Similar to their behavior in solution, the Φ_{PL} of the thin films are higher under a nitrogen atmosphere than when exposed to air, confirming the presence of TADF in solid state. Similar to the solution-state emission spectra, the thin film emission spectra of the four oxadiazole emitters were blue-shifted compared to that of **2CzPN**. Figure 2d shows the corresponding PL decay curves. For each of the emitters, there is a fast prompt decay ($\tau_{\text{p}} \approx 10 \text{ ns}$) and several very long delayed emission components. The dominant delayed components, τ_{d} , were estimated to be $\tau_{\text{d1}} = 3.0 \text{ ms}$ (75%, amplitude in multi-exponential fit to data) and $\tau_{\text{d2}} = 40 \text{ ms}$ (19%) for compounds **2CzdOXDMe**, $\tau_{\text{d1}} = 4.9 \text{ ms}$ (63%) and $\tau_{\text{d2}} = 45.2 \text{ ms}$ (27%) for **2CzdOXDPh**, $\tau_{\text{d1}} = 2.3 \text{ ms}$ (77%) and $\tau_{\text{d2}} = 23.2 \text{ ms}$ (19%) for **2CzdOXD4CF₃Ph**, and $\tau_{\text{d1}} = 1.5 \text{ ms}$ (83%) and $\tau_{\text{d2}} = 17 \text{ ms}$ (10%) for **2CzdOXD4MeOPh**.

For device applications, 1,3-bis(carbazol-9-yl)benzene (mCP) and 2,8-bis(diphenylphosphoryl) dibenzo[b,d] thiophene (PPT) hosts were selected, due to their hole or electron transporting ability, respectively, in combination with the high triplet energy ($E_T = 2.9$ eV and 3.0 eV for mCP⁷⁴ and PPT⁷⁵, respectively). Figure 3 and Table 4 summarize the photophysical properties of the emitters embedded in vacuum-deposited mCP or PPT matrices. Due to the high singlet energies of these matrices, Förster energy back-transfer cannot occur and the emission of the doped films originates solely from the dopant (Figure S42). The Φ_{PL} values of the emitters embedded in mCP or PPT remain relatively high, ranging from 14 to 55%. They are, however, lower compared to the Φ_{PL} recorded in PMMA films. The singlet-triplet gap, ΔE_{ST} , of each of the emitters was estimated from the difference in the onset of the fluorescence and phosphorescence spectra recorded at 77K (Figure 3a). Overall, the compounds exhibit experimentally determined ΔE_{ST} that are comparable to 2CzPN,⁷⁶ and range from 0.30 to 0.46 eV (Figure 3b), which are in line with the theoretical calculations (*vide infra*). All of the compounds showed prompt and delayed emission in the vacuum-deposited films (Figure 3c). The spectroscopic evidence signifies that the four oxadiazole compounds are indeed TADF emitters.

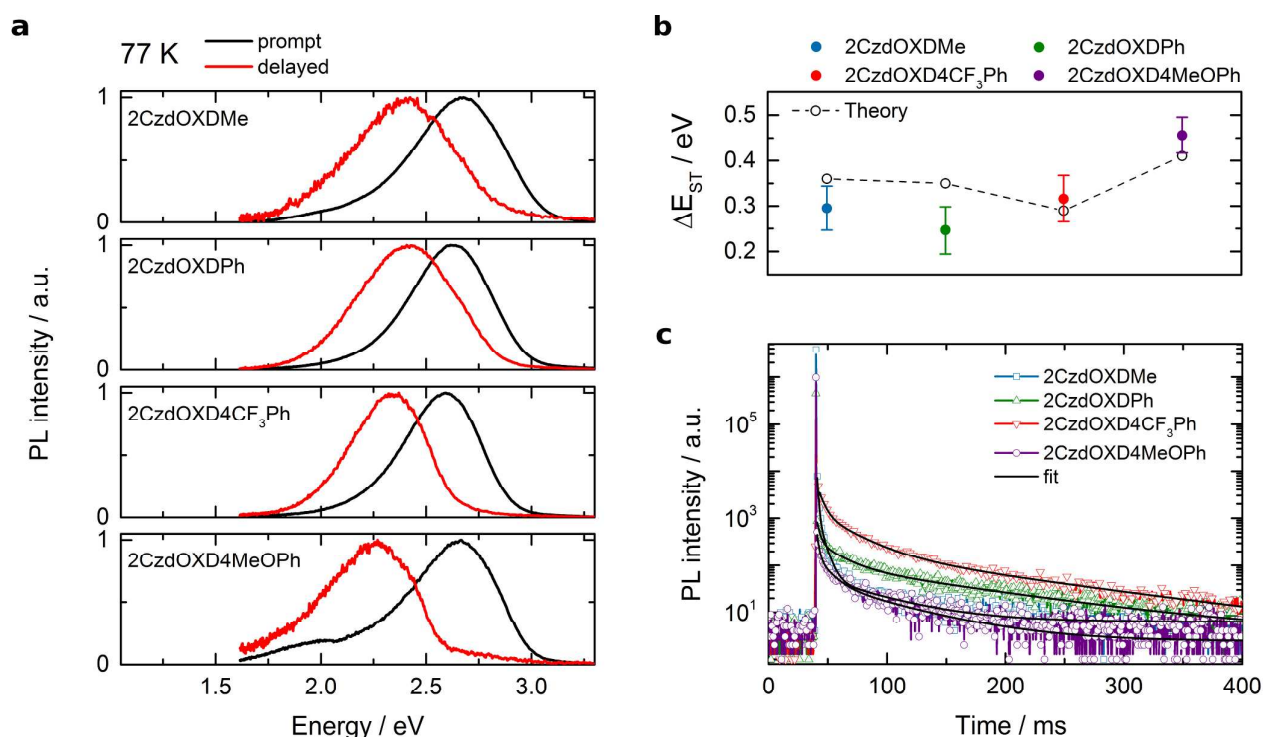


Figure 3. a) Prompt and delayed spectra of emitters embedded in mCP (2CzdOXDMe and 2CzdOXDMeOPh) or PPT (2CzdOXDPh and 2CzdOXD4CF₃PH) hosts at 6 wt%. Measurements are performed at 77 K. b) Summary of the ΔE_{ST} values estimated from the onset of fluorescence and phosphorescence spectra and compared to the DFT calculations. c) Corresponding PL decay curves at room temperature ($\lambda_{exc} = 378$ nm).

Table 4. Summary of thin film photophysical properties of studied compounds embedded in mCP or PPT hosts.

Film	λ_{PL}	FWHM	Φ_{PL}^a / %	τ_{PL} / ns	Prompt:delayed	ΔE_{ST} / eV
	/	/ nm			/ %	
	nm					
mCP:2CzdOXDMe	435	73	29.3	5.7, $1 \cdot 10^6$	92.0:8.0	
			(27.0)	(86%),		0.30 ± 0.05
				$4.8 \cdot 10^6$		

					(13%)	
PPT:2CzdOXDPh	454	85	43.9	9.5,	88.2:11.8	
			(38.7)	2.4·10 ⁶		0.25±0.05
				(62%),		
				17.8·10 ⁶		
				(27%)		
PPT: 2CzdOXD4CF ₃ Ph	474	83	54.9	6.0,	74.6:25.4	
			(41.0)	4.5·10 ⁶		0.32±0.05
				(67%),		
				25.8·10 ⁶		
				(28%),		
mCP:2CzdOXD4MeOPh	449	77	14.9	8.1,	95.1:4.9	0.46±0.04
			(14.1)	1.7·10 ⁶		
				(69%),		
				10.1·10 ⁶		
				(23%),		

^a Values in parentheses are for aerated samples.

Theoretical Calculations

Further characterization of the optoelectronic properties of these compounds has been carried on with the help of DFT calculations. To do so, we used the methodology developed by Moral *et al.* offering a particularly accurate description of the electronic structure of materials for OLED applications.⁷⁷ Not surprisingly, the HOMO energies of the different compounds remain almost constant as in each case the HOMO is localized on the carbazole moiety (Figure 4). These are ca. 0.1 eV more stabilized than the HOMO for 2CzPN, indicating that the nature of the acceptor moiety modestly modulates the HOMO. The LUMO

energies on the other hand decrease significantly, showing that the synthetic approach allows for near-independent tuning of the frontier orbital energies upon replacing cyano groups in 2CzPN with oxadiazole acceptors. Not surprisingly, the replacement of cyano groups with oxadiazoles results in a more delocalized LUMO for the four oxadiazole derivatives while the HOMO remains mainly localized on the carbazole groups and the central phenyl ring (Figure 4).

Expectedly, because of the lower electron-withdrawing character of methyl-substituted oxadiazole, **2CzdOXDMe** possesses the highest energy LUMO in the series, more than 0.4 eV destabilized compared to **2CzPN**, resulting in a significantly bluer emission (see S_1 energy in Table 5). Replacing the methyl group with phenyl as is the case in **2CzdOXDPh** results in a more modest 0.24 eV destabilization of the LUMO compared to **2CzPN**. Interestingly, by grafting CF_3 -substituted phenyl groups to the oxadiazole units, the LUMO energy of **2CzdOXD4CF₃Ph** is stabilized with respect to **2CzdOXDPh** and in fact is close to that of **2CzPN**; conversely the LUMO energy of **2CzdOXD4MeOPh** is destabilized relative to **2CzdOXDPh**.

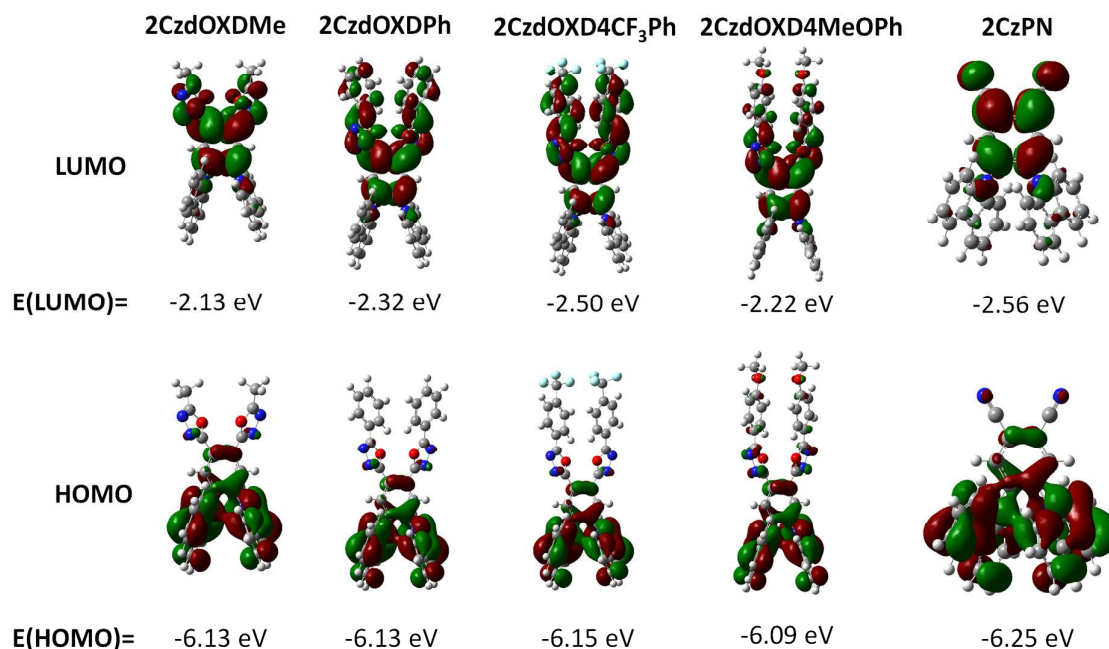


Figure 4. Frontier orbitals contour plots and energies of the different derivatives as well as **2CzPN** considering acetonitrile as a solvent.

The calculations are able to predict the emission colors of both **2CzPN** and the four oxadiazole derivatives. Specifically, the S_1 energies of **2CzdOXDMe** appears as the bluest while **2CzdOXD4CF₃Ph** is slightly red-shifted compared to **2CzPN**. All molecules exhibit similar electronic structure with T_2 lying in between T_1 and S_1 .

We note that an additional T_3 excited state lies below S_1 in **2CzdOXD4MeOPh**. Considering the electronic structure, the up-conversion to S_1 could potentially arise either directly from T_1 or from an intermediate triplet excited state after reverse internal conversion. The T_1 and S_1 excited states are mainly characterized by HOMO to LUMO transitions with a pronounced intramolecular charge-transfer (ICT) character. Interestingly, the singlet-triplet splitting, ΔE_{ST} , moderately decreases when going from **2CzdOXDMe** to **2CzdOXD4CF₃Ph**, in line with a slight decrease of the oscillator strength. We further characterize the degree of

spatial separation between occupied (ϕ_i) and virtual (ϕ_a) molecular orbitals, Δr , relative to a particular excited state by evaluating the averaged hole-electron distance based on the computation of the detachment and attachment centroids radius vectors and the calculation of the distance between them using the NANCY_EX package.⁷⁸⁻⁸⁰ We mention that this approach appears to be perfectly equivalent⁷⁸ to the one we used previously and which is based on the corresponding centroid of the Natural Transition Orbitals involved in the studied excited-state transition.⁷⁷

A cutoff $\Delta r > 1.5\text{--}2.0$ Å has been proposed previously to both distinguish, and characterize, a charge-transfer excitation,⁸¹ and it was shown that the magnitude of the ΔE_{ST} is inversely proportional to Δr .⁷⁷ Recently, it has been demonstrated that in order to obtain TADF emitters with very small ΔE_{ST} , both Δr relative to the excitation from S_0 to T_1 [$\Delta r(T_1)$] and S_1 [$\Delta r(S_1)$] should be large, and in particular the triplet component was found to play a dominant role in determining ΔE_{ST} of the different compounds studied.⁸² When examining Δr in Table 5, all Δr reported confirm the expected ICT character that could be inferred from the orbital localization (Figure 4). Among all the compounds, **2CzPN** and **2CzdOXD4CF₃Ph** show the largest $\Delta r(T_1)$ and exhibit the smallest ΔE_{ST} while **2CzdOXD4MeOPh** shows the largest ΔE_{ST} and smallest $\Delta r(T_1)$, in agreement with what was proposed by Huang and co-workers.⁸²

Table 5. Excitation energies, ΔE_{ST} , oscillator strengths and compositions of the excited states in terms of monoelectronic transitions (molecular orbitals) for **2CzPN** and its derivatives.

Compound	States	Energy	O.S.	Main MO component of the	Δr
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		(eV)	(a.u.)	transitions from S ₀ (%)	(Å)
2CzdOXDMe	T ₁	2.88	-	HOMO → LUMO (80.9%)	2.40
	S ₁	3.24	0.13	HOMO → LUMO (97.9%)	3.82
	ΔE(S₁-T₁)	0.36			-
2CzdOXDPh	T ₁	2.75	-	HOMO → LUMO (75.7%)	2.43
	S ₁	3.10	0.12	HOMO → LUMO (97%)	4.37
	ΔE(S₁-T₁)	0.35			-
2CzdOXD4CF₃Ph	T ₁	2.68	-	HOMO → LUMO (76.9%)	3.12
	S ₁	2.97	0.10	HOMO → LUMO (95.8%)	4.91
	ΔE(S₁-T₁)	0.29			-
2CzdOXD4MeOPh	T ₁	2.76	-	HOMO → LUMO (71.4%)	1.24
	S ₁	3.17	0.13	HOMO → LUMO (97.2%)	3.73
	ΔE(S₁-T₁)	0.41			
2CzPN	T ₁	2.67	-	HOMO → LUMO (87.7%)	2.80
	S ₁	2.98	0.11	HOMO → LUMO (96.5%)	3.70
	ΔE(S₁-T₁)	0.31			-

Crystal structures

Suitable crystals for X-ray analysis were obtained for all of **2CzdOXDMe**, **2CzdOXDPh**, **2CzdOXD4CF₃Ph** and **2CzdOXD4MeOPh**, as well as for the reference compound **2CzPN** (Figure 5). Crystals of **2CzdOXDMe** and **2CzPN** were grown by the slow evaporation of mixed solutions of toluene/hexane, crystals of **2CzdOXD4CF₃Ph** by the slow evaporation of mixed solutions of toluene/MeCN, while crystals of **2CzdOXDPh** and **2CzdOXD4MeOPh** were grown by the vapour diffusion of Et₂O into concentrated solutions of 1,2-dichloroethane (**2CzdOXDPh**) or dichloromethane (**2CzdOXD4MeOPh**). All four new compounds exhibit similar conformations, with both carbazole rings disposed in a highly twisted arrangement with respect to the plane of the central benzene [dihedral angles of 58.51(3)–67.64(7)°]. In comparison to the structure of **2CzPN**,⁸³ the carbazole units in **2CzdOXDMe**, **2CzdOXDPh**, **2CzdOXD4CF₃Ph** and **2CzdOXD4MeOPh** are closer to orthogonal to the central benzene [**2CzPN** dihedral angles of 50.60(4)–60.83(4)°]. The oxadiazole heterocycles in the four

emitters are less restrained in the orientations they can occupy due to their smaller size, and show a much wider range of dihedral angles relative to the central benzene than the carbazole fragments, with a general pattern of having one oxadiazole ring close to coplanar with the benzene, and one close to orthogonal [dihedral angles 6.18(8)–21.77(10) for more coplanar rings, and 49.32(5)–78.11(11)° for more orthogonal]. In **2CzdOXDPh**, **2CzdOXD4CF₃Ph** and **2CzdOXD4MeOPh**, the oxadiazole moiety and its aryl substituent are approximately coplanar [dihedral angles of 4.04(9)–28.27(6)°], indicative of a degree of conjugation between these ring systems in the solid-state structures. Additionally, one further distinction is seen in the structures. For **2CzdOXDMe** and **2CzdOXDPh**, the oxadiazoles are oriented such that they are aligned closer to parallel than antiparallel, whereas for **2CzdOXD4CF₃Ph** and **2CzdOXD4MeOPh**, the converse is the case.

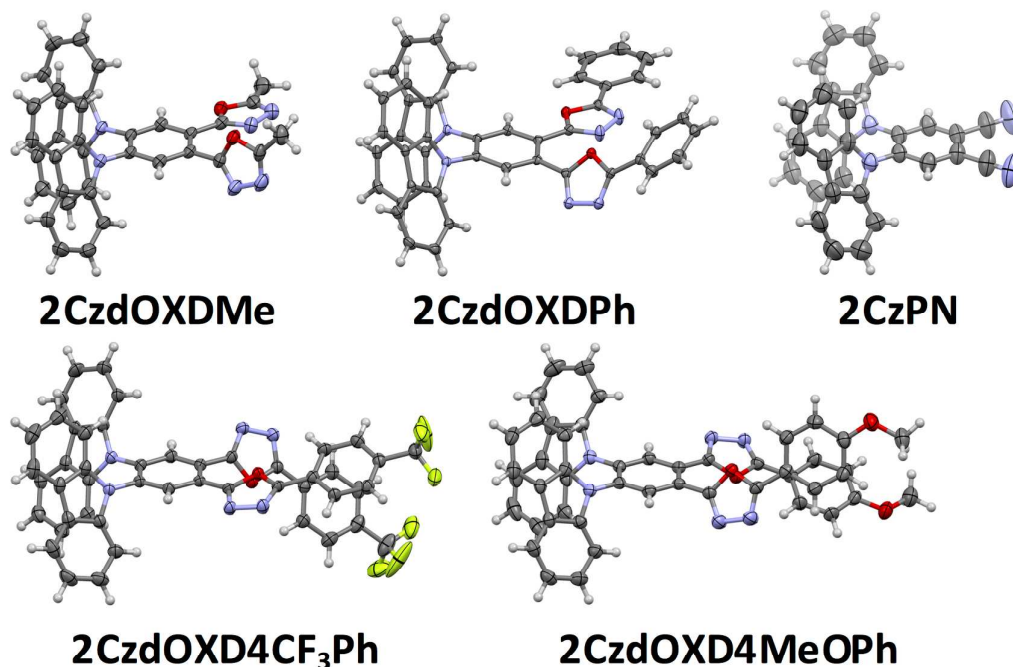


Figure 5. Structures of **2CzdOXDMe**, **2CzdOXDPh**, **2CzdOXD4CF₃Ph**, **2CzdOXD4MeOPh**, and **2CzPN** (50% probability ellipsoids; co-crystallized solvent molecules omitted). Heteroatoms: N, light blue; F, green; and O, red.

Organic Light-Emitting Diodes

The four oxadiazole emitters were then integrated into OLED devices. Figure 6 shows the architecture employed to build sky-blue to deep-blue OLEDs. The general device structure consisted of indium tin oxide ITO (90 nm)/NPB (35 nm)/mCP (10 nm)/EML (15 nm)/PPT (10 nm)/TmPyPB (30 nm)/LiF (1 nm)/Al (100 nm). NPB is *N,N'*-bis(naphthalen-1-yl)-*N,N'*-bis(phenyl)-benzidine and acts as the hole transport layer. TmPyPB is 1,3,5-tri(*m*-pyrid-3-yl-phenyl)benzene and acts as the electron transporting layer (ETL). Devices **1-5** comprising different emission layers (EMLs) incorporating the oxadiazole emitters were built, and these were compared to a reference Device **R1**, based on **2CzPN**. The Device **R1** was based on the device stack reported by Adachi and co-workers:⁷⁶ ITO (90 nm)/NPB (35 nm)/mCP (10 nm)/ **2CzPN**:mCP (6 wt%, 15nm)/PPT (10 nm)/TPBi (40 nm)/LiF (1 nm)/Al (100 nm). Here, the ETL used was TPBi, which is 1,3,5-tris(1-phenyl-1H-benzimidazol- 2-yl)benzene. The thin layer of 10 nm mCP (PPT) before (after) the EML in all of the devices acts as the electron (hole) blocking layer.

The EML of Device **1** consisted of **2CzdOXDMe**:mCP (6 wt%). Devices **2** and **3** comprised **2CzdOXDPh**:PPT (6 wt%) and **2CzdOXD4CF₃Ph**:PPT (6 wt%) as their EMLs. For **2CzdOXD4MeOPh**, devices employing both mCP (Device **4**) and PPT (Device **5**) as hosts were fabricated. The reason for using mCP for **2CzdOXDMe** is that the LUMO level of **2CzdOXDMe** (-2.7 eV) matches the LUMO of PPT,⁷⁶ and thus efficient electron trapping by

the dopant would not be achieved in a PPT matrix. On the other hand, mCP is predominantly hole conductive^{74,84} and hole injection and conduction is more efficient compared to electron transport in the mCP device. Therefore, better charge balance within the EML was expected when using PPT as a host and thus PPT was used as a matrix material for emitters **2CzdOXDPh** and **2CzdOXD4CF₃Ph**. Due to the intermediate LUMO value of **2CzdOXD4MeOPh** (-2.74 eV), it was tested in both mCP and PPT.

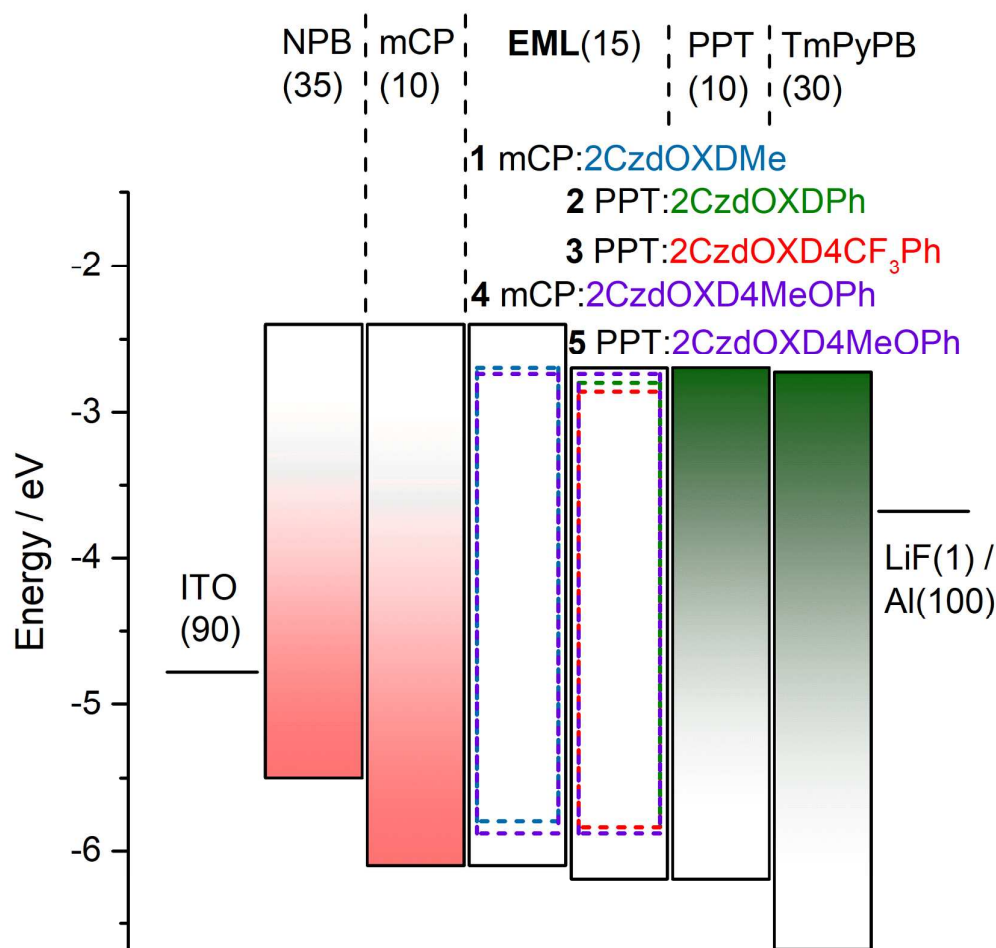


Figure 6. Functional layer sequence of the devices **1-5** comprising oxadiazole emitters.

Figure

Figure 7 and Table 6 summarise the OLED performance. The electroluminescence (EL) spectra shift systematically from sky-blue emission exhibited by Device **R1** and Device **3** ($\lambda_{\text{EL}} = 480$ nm and 474 nm, respectively) to deep-blue emission exhibited by Devices **1, 2,**

4 and 5, with corresponding peak emission at $\lambda_{\text{EL}} = 446$ nm, 455 nm, 446 nm and 448 nm, respectively (Figure 7a). In agreement with photophysical data (Tables 3 and 4), devices fabricated with **2CzdOXD4CF₃Ph** (Device 3) demonstrated the most red-shifted electroluminescence compared with the other three oxadiazole emitters due to the presence of the electron-withdrawing CF₃ groups, which result in **2CzdOXD4CF₃Ph** possessing the deepest LUMO (-2.86 eV, Table 2) among all oxadiazole emitters. In general, the electroluminescence spectra of the oxadiazole emitters agree very well with photoluminescence spectra obtained from their thin films doped in mCP or PPT hosts (Table 4) and the trend of their emission energies can be correlated to their corresponding LUMO levels (Table 2). Devices 2 and 3 show the highest turn-on voltages V_{on} of 4.30 V and 4.60 V, respectively, and flat current-voltage characteristic (Figure 7b). This could be indicative of high series resistance due to poor charge injection and transport when using **2CzdOXDPh** and **2CzdOXD4CF₃Ph** doped PPT system as the EMLs. On the other hand, Devices 1, 4 and 5, show steep current-voltage characteristics (Figure 7b) and V_{on} of 3.85 V, 4.15 V and 4.05 V, respectively.

Figure 7c shows the external quantum efficiency (EQE) versus the current density of the OLEDs. Device 3 reaches a maximum EQE_{max} of 11.2% at low brightness, indicating the presence of an efficient triplet harvesting via the TADF mechanism (Figure 7c). However, the EQE of 3 drops by almost a factor of five to 2.5% at 100 cd m⁻². Similar trends of relatively high EQE at low brightness (4.7%, 6.8%, 6.6% and 4.2% for Devices 1, 2, 4, and 5, respectively) and a strong efficiency roll-off are also observed for the other devices. Such efficiency roll-off might be caused by the bimolecular annihilation reactions due to the presence of the long-lived triplet states (τ_{PL} ranging from 1.0 to 25.8 ms), as evidenced from the photophysical studies of the thin films of oxadiazole emitters doped in mCP or PPT hosts

(Table 4). The most severe efficiency roll-off was observed for Devices **2** and **3**, comprising emitters **2CzdOXDPh** and **2CzdOXD4CF₃Ph**, respectively, with the corresponding critical current densities j_{crit} in the range of 0.05 mA cm^{-2} . Here, j_{crit} denotes the current density at which the EQE drops to half of its maximum value. In comparison, the lowest roll-off was observed for the device with **2CzdOXDMe** as the emitter, device **1** ($j_{\text{crit}} = 13 \text{ mA cm}^{-2}$). The average delayed lifetime (τ_{avg}) of emitters **2CzdOXDPh** and **2CzdOXD4CF₃Ph** and **2CzdOXDMe** is indicative of the triplet state lifetime in these compounds and is estimated to be 6.3 ms, 10.2 ms and 1.5 ms, respectively (Table 4). The shorter τ_{avg} leads to the less pronounced efficiency roll-off, in line with the bimolecular annihilation pathway being the limiting factor for efficient OLED operation at high current density and brightness. Thus, further work on optimising exciton dynamics in the EML and minimizing bimolecular annihilation reactions is needed. Additionally, it is important to note that the reference device **R1** used in this work does not represent the optimal device structure for the sky-blue emitter **2CzPN**. Indeed, the selection of the functional layer sequence is crucial to minimize the electrical losses in blue TADF OLEDs. For example, Zhang *et al.*⁸⁵ showed that replacing TPBi with diphenylphosphine oxide-based ETLs that exhibit high electron mobility and high E_{T} can enhance the **2CzPN**-based OLED performance up to $\text{EQE}_{\text{max}} = 17.4\%$. Replacement of mCP as the EBL with a higher hole mobility and thermal stability material 3,5-di(9H-carbazol-9-yl)-*N,N*-diphenylaniline (DCDPA)⁸⁶ has been shown to increase the efficiency further to $\text{EQE}_{\text{max}} = 19.2\%$.⁸⁷ Finally, Sun *et al.*⁸⁸ fabricated OLEDs comprising a mixed host layer for the TADF dopant to boost the performance further to 21.8%, which was also shown to be the efficiency limit for **2CzPN**-based devices. Similar improvements are thus expected to enhance device performance when oxadiazole emitters are used. Nevertheless, this work demonstrates that a systematic shift toward deep-blue electroluminescence is possible by replacing the CN accepting groups in **2CzPN** by oxadiazole unit containing moieties. Figure

7d shows the corresponding color coordinates of Devices **1-5** in comparison to the reference Device **R1**. Devices **1**, **2**, **4**, and **5** all exhibit CIE x,y coordinates < 0.2 and, thus, are well suited as deep blue emitters.

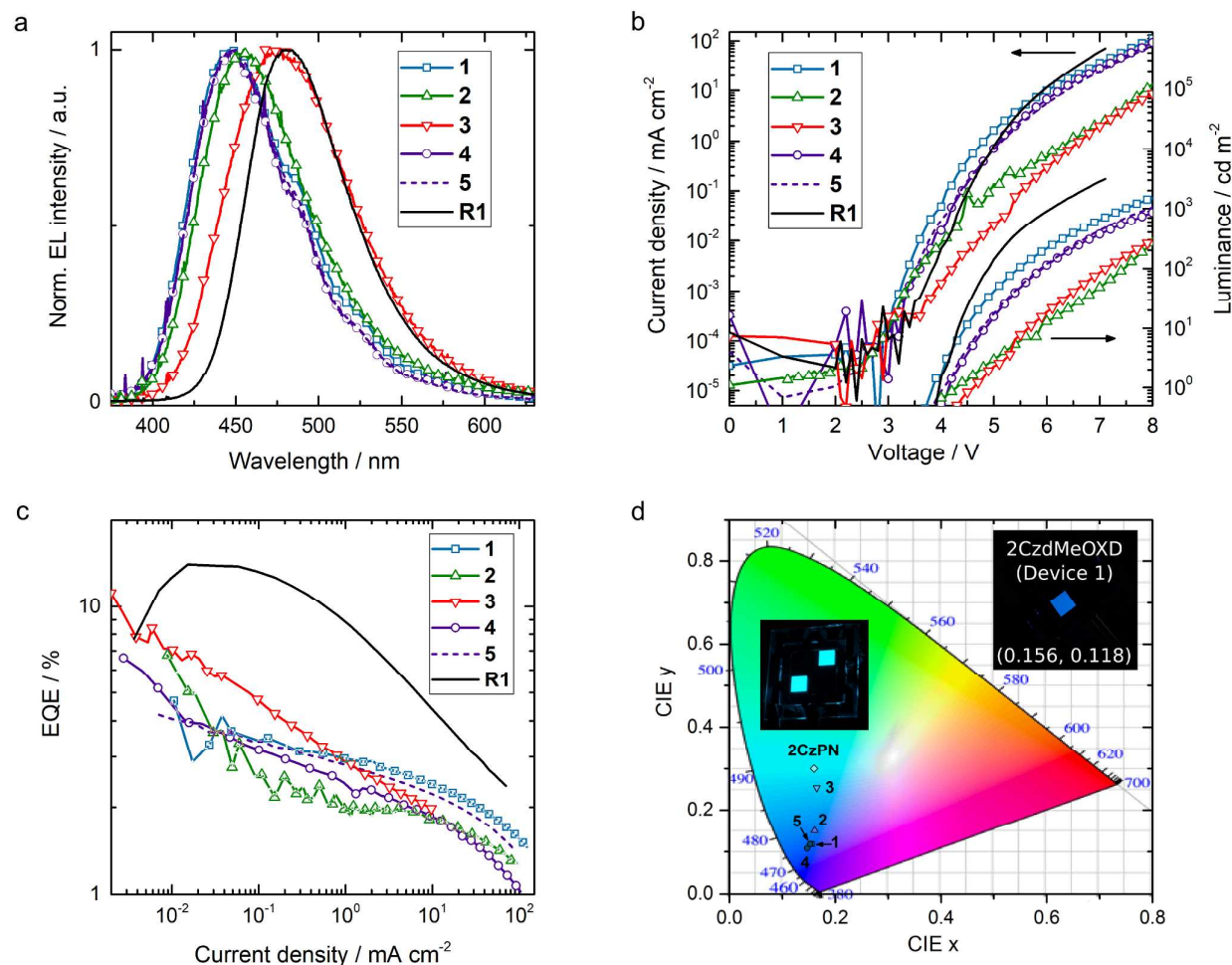


Figure 7. OLED performance of devices based on sky-blue to deep-blue emitters. **a)** Electroluminescence spectra; **b)** current density-voltage-luminance characteristics; **c)** external quantum efficiency dependence on current density; **d)** CIE color coordinates of devices **1-5** and Device **R1** based on **2CzPN** for comparison. Inset shows photographs of Device **1** and Device **R1**.

Table 6. Summary of OLED performances.

Emitter	Host	$V_{\text{on}} / \text{V}^{\text{a}}$	$\lambda_{\text{EL}} / \text{nm}$	FWHM / nm	CIE (x, y)	$\text{EQE}_{\text{max}} / \text{EQE}_{100} / \text{EQE}_{1000} / \%$ ^b	$\text{PE}_{\text{max}} / \text{lm W}^{-1}$	$\text{CE}_{\text{max}} / \text{cd A}^{-1}$
2CzdOX DMe	mCP	3.85	446	78	(0.156, 0.118)	4.7 / 2.6 / 1.8	3.7	4.2
2CzdO XDPH	PPT	4.30	455	75	(0.161, 0.150)	6.8 / 2.0 / 1.4	5.9	7.5
2CzdO	PPT	4.60	474	78	(0.165, 0.254)	11.2 / 2.5 / - ^c	14.5	18.9

XD4CF ₃ Ph								
2CzdO	mCP	4.15	446	73	(0.147, 0.108)	6.6 / 2.0 / 0.8	5.0	5.7
XD4Me								
OPh								
2CzdO	PPT	4.05	448	73	(0.153, 0.118)	4.2 / 2.4 / 1.5	3.2	3.8
XD4Me								
OPh								
2CzPN	mCP	3.95	480	71	(0.160, 0.300)	14.0 / 10.3 / 4.0	18.4	25.9

^a. Defined as the lowest operating voltage at the luminance of >1 cd m⁻²; ^b. Max = Maximum value, 100 = measured at 100 cd m⁻², 1000 = measured at 1000 cd m⁻²; ^c. 1000 cd m⁻² not reached.

Conclusions

This study shows the viability of using oxadiazole acceptor units to blue-shift the emission while conserving the TADF properties of the emitters in the solid-state. Compared to **2CzPN**, the replacement of the nitrile acceptors for oxadiazole units destabilized the LUMO level while keeping the HOMO essentially unchanged, which was observed both experimentally and corroborated by DFT calculations, causing an increase in the emission energy. The TADF nature of the compounds was confirmed by a combination of oxygen-dependence measurements on the photoluminescence quantum yield, time-resolved measurements clearly showing prompt and delayed components in the film. The experimentally measured and calculated ΔE_{ST} are in strong agreement. OLED devices using **2CzdOXD4MeOPh** produced deep blue light with CIE coordinates of (0.147, 0.108) and maximum external quantum efficiency of 6.6% while a more efficient (EQE_{max} of 11.2%) sky-blue device was fabricated (0.165, 0.254) with **2CzdOXD4CF₃Ph**. The efficiencies obtained are demonstrative of TADF being operational in the OLED devices. In future, the observed strong roll-off of the devices needs to be addressed using improved device design.

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Supporting Information. Complete experimental protocol. NMR spectra. HR-mass spectra. HPLC traces. CIF files. (CCDC: 1852947-1852950, 1436867) for **2CzdOXDMe**, **2CzdOXDPh**, **2CzdOXD4CF₃Ph**, **2CzdOXD4MeOPh**, and **2CzPN**. Supplementary photophysical data. Supplementary computations and coordinates.

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